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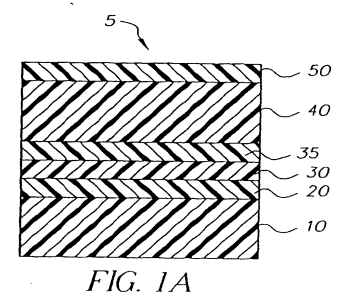
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(54) Compensator layer and liquid crystal cell with such a layer

(57) Disclosed is a liquid crystal cell comprising a liquid crystal layer enclosed between two constraints and having contiguous to a surface of one of the constraints one or more compensator layers, each contain-

ing a transparent amorphous polymeric birefringent material having an out-of plane birefringence more negative than -0.005. The invention also provides a liquid crystal display.



Description

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[0001] The present invention relates to a liquid crystal cell having contiguous to an exterior surface of a constraint thereof a compensator layer containing an amorphous polymeric birefringent material having an out-of plane birefringence more negative than -0.005. The invention also provides an LC display and a process for making such a cell.

[0002] Triacetylcellulose (TAC, also called cellulose triacetate) film has traditionally been used by the photographic industry due to its unique physical properties and flame retardance. TAC film is also the preferred polymer film for use as a cover sheet for the polarizers used in liquid crystal displays. It is the preferred material for this use because of its extremely low in-plane birefringence. Its out of plane birefringence is also small (but not zero), and is useful in providing some optical compensation to the LCD.

[0003] Intrinsic birefringence describes the fundamental orientation of a material at a molecular level. It is directly related to the molecular structure (bond angles, rotational freedom, presence of aromatic groups, etc.) of the material. The intrinsic birefringence is not affected by process conditions (temperature, stresses, pressures) used to make a macroscopic object.

[0004] Crystalline and liquid crystalline materials have the convenient property that their intrinsic birefringence manifests itself almost perfectly when they are assembled into a macroscopic article. Layers of crystalline and liquid crystalline molecules often can be manufactured such that all the molecules in the article are in registry with each other and thus preserve their fundamental orientation. The same is <u>not</u> true when making layers of an amorphous polymeric material. Their intrinsic birefringence can be highly modified by the manufacturing process. Thus, the measured birefringence of an actual article will be a resultant of its intrinsic birefringence and the manufacturing process. Because we are dealing with such amorphous polymeric materials, the following definitions refer to this measured birefringence and not intrinsic birefringence.

[0005] In-plane birefringence means the difference between n_x and n_y , where x and y lie in the plane of the layer. n_x will be defined as being parallel to the casting direction of the polymer, and n_y being perpendicular to the casting direction of the polymer film. The sign convention used will be $n_x - n_y$.

[0006] Out of-plane birefringence means the difference between n_z and the average of n_x and n_y , where x and y lie in the plane of the layer and z lies in the plane normal to the layer. The sign convention used will be: $n_z = [(n_x + n_y)/2]$. TAC typically has a negative out of plane birefringence as its n_z is smaller than its n_x and n_y .

[0007] In-plane retardation (Re) means the product of in-plane birefringence and layer thickness (t). Thus Re = t (n_x - n_y)

[0008] Out-of-plane retardation (Rth) means the product of out-of-plane birefringence and layer thickness (t). Thus Rth = t $(n_z - [(n_x + n_y)/2])$.

[0009] Synthetic polymer films (such as polycarbonate or polysulfone) are often used to enhance the minimal optical compensation that TAC provides. These synthetic polymers films are attached to the rest of the display by adhesive lamination.

[0010] Generally in the field of optical materials, the synthetic polymer film is used as an optically anisotropic film (having a high retardation value), while a TAC film is used as an optical isotropic film (having a low retardation value). [0011] Japanese Published Patent Application JP1999-95208 describes a liquid crystal display having an optical compensator (having high retardation) prepared by uniaxial stretching of a high polymer film. Such polymers include polyesters, polycarbonate or polysulfone. This stretching step is essential to obtain the desired optical properties. This stretching affects both in- and out-of-plane retardation simultaneously. These two orthogonal retardations cannot be independently controlled by this method. Also, producing uniform optical compensators by this method is described as being difficult.

[0012] This application also describes a compensator where the inventor uses an exfoliated inorganic clay material in a polymeric binder coated on top of a TAC support. The exfoliated inorganic clay material in this layer is the optically active material, not the polymeric binder.

[0013] World patent WO 01/31394 A2 discusses the use of the color filter array layer as a source of additional out-of-plane retardation for a liquid crystal display. The color filter array is located within the constraints of the liquid crystal cell. The use of an aromatic polyimide binder rather than a polyacrylate binder for the color filter array dyes provides enhanced retardation. The overall retardation is achieved with the combination of the color filter array retarder plus optional additional out-of-plane retardation from the TAC used as a supporting member for the polarizers.

[0014] The proposal to select the binder for the color filter array with retardation in mind has an advantage versus polarizer-based retarders that are laminated to the liquid crystal cell: mechanical stresses to the display induced by room condition changes or perhaps direct shock can cause polarizer-based retarders to move relative to the liquid crystal cell. Retarders coated directly on the glass substrate are more rigidly held in registry with the cell, and thus do not suffer this problem. However the requirement that this color filter array be also a retarder means that this layer must serve two purposes: color filtering and adding retardation. This limits the potential thickness to be considered for this layer. This layer must also be pixilated, and this adds additional complications. Finally it is taught on the internal

surface of the constraint only, where the color filter array is located.

[0015] It is a problem to be solved to provide a liquid crystal cell that is readily manufactured and that readily provides the required degree of in-plane and out-of-plane compensation while reducing the problems associated with a laminated compensator.

[0016] The invention provides liquid crystal cell having contiguous to a surface of a constraint thereof one or more compensator layers, each containing a transparent amorphous polymeric birefringent material having an out-of plane birefringence more negative than -0.005. The invention also provides a liquid crystal display and a process for preparing a compensator of the invention.

[0017] The invention cell is readily manufactured and provides the required degree of in-plane and out-of-plane compensation.

FIG. 1A is a cross-sectional schematic of an embodiment of the invention with the amorphous polymeric compensator layer on the side of the constraint opposite the liquid crystal

FIG. 1B is a cross-sectional schematic of an embodiment of the invention with the amorphous polymeric compensator layer on the side of the constraint adjacent to the liquid crystal

FIG. 2A is an exploded view schematic of a liquid crystal display with one amorphous polymeric compensator layer of the invention

FIG. 2B is an exploded view schematic of a liquid crystal display with two amorphous polymeric compensator layers of the invention

[0018] The invention is summarized above.

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[0019] The present invention is not limited by the requirements of WO 01/31394 mentioned above.

[0020] The present invention provides a liquid crystal cell having contiguous to at least one surface of a constraint thereof a compensator layer containing a birefringent amorphous polymeric material having an out-of plane birefringence more negative than -0.005.

[0021] As used herein, constaints are the two principal supporting members of the LC cell (typically glass) that sandwich the switchable liquid crystal layer (and typically the color filter array, black matrix, and thin film transistor, alignment and electrode layers, and other optional layers) and are normally at least 10 micrometers thick. The term "transparent" is used in its normal sense to mean a layer that absorbs little or no visible light.

[0022] The liquid crystal cell referred to herein extends from the external surface of one constraint to the other, including any compensator layer on the constraint surfaces.

[0023] Amorphous polymeric materials are used for this optical compensator. In this case amorphous means that the optical compensator would not produce any sharp diffraction peaks when exposed to X-ray diffraction analysis. Crystalline polymers. liquid crystal molecules and crystalline inorganic materials would produce such sharp peaks when subjected to such X-ray diffraction analysis. Such amorphous materials are desirably suitable to be solvent cast or coated such as TAC, polycarbonates, cyclic polyolefins, polyesters, and polyimides.

[0024] Typical lamination uses pressure sensitive adhesive layers of greater than 4 micrometers in thickness. As used herein, the term contiguous means without the use of any intervening laminating adhesive layer and therefore contemplates the possible use of only a very thin (0.2µm or less) adhesion promoting layer or an adhesion promoting surface treatment such as corona discharge, plasma glow discharge, or flame treatment. Other adhesion enhancing methods could be employed as known to those skilled in the art.

[0025] The compensator layer will typically be solvent coated onto the constraint exterior surface. This solvent coating could be accomplished by spin coating, hopper coating, gravure coating, wire bar coating, spray coating, or other coating methods known to those skilled in the art.

[0026] The compensator layer is coated from a solution containing a polymer that yields high negative birefringence upon solvent coating. To produce negative birefringence (negative retardation), polymers that contain non-visible chromophore groups such as vinyl, carbonyl, amide, imide, ester, carbonate, sulfone, azo, and aromatic groups (i.e. benzene, naphthalate, biphenyl, bisphenol A) in the polymer backbone will be used, such as polyesters, polycarbonates, polyimides, polyetherimides, and polythiophenes. One could also add fillers and non-polymeric molecules to these polymers forthis contiguous layer.

[0027] Examples of suitable polymers for use in the second layer are 1)poly(4,4'-hexafluoroisopropylidene-bisphenol) terephthalate-co-isophthalate, 2)poly(4,4'-hexahydro-4,7-methanoindan-5-ylidene bisphenol) terephthalate, 3) poly(4,4'-isopropylidene-2,2'6,6'-tetrachlorobisphenol) terephthalate-co-isophthalate, 4) poly(4,4'-hexafluoroisopropylidene)-bisphenol-co-(2-norbornylidene)-bisphenol terephthalate, 5) poly(4,4'-hexahydro-4,7-methanoindan-5-ylidene)-bisphenol-co-(4,4'-isopropylidene-2,2',6,6'-tetrabromo)-bisphenol terephthalate, and 6) poly(4,4'-isopropylidene-bisphenol-co-4,4'-(2-norbornylidene) bisphenol) terephthalate-co-isophthalate or copolymers of any of the foregoing.

[0028] A chromophore is defined as an atom or group of atoms that serve as a unit in light adsorption. (Modern

Molecular Photochemistry Nicholas J. Turro Editor, Benjamin/Cummings Publishing Co., Menlo Park, CA (1978) Pg 77). Typical chromophore groups include vinyl, carbonyl, amide, imide, ester, carbonate, aromatic (i.e. phenyl, naphthyl, biphenyl, thiophene, bisphenol), sulfone, and azo or combinations of these chromophores. A non-visible chromophore is one that has an absorption maximum outside the range of 400-700nm.

[0029] Desirably, polymers to be used in the compensator layer will not have chromophores off of the backbone. An example of such an undesirable polymer with chromophores in and off the backbone would be polyarylates possessing significant proportions of the fluorene group.

[0030] The glass transition temperature (Tg) of the polymers used in the compensator layer is significant. Tg values above 180° C are desirable to achieve preferred results.

[0031] The polymers used in the contiguous compensator layer can be synthesized by a variety of techniques: condensation, addition, anionic, cationic or other common methods of synthesis can be employed.

[0032] The thickness of this layer is usually less than 30μm. Typically it is from 0.1μm to 20μm. Conveniently it is from 1.0μm to 10μm and desirably from 2μm to 8μm.

[0033] The compensator layer is suitably of sufficient thickness so that the out-of-plane retardation of the second layer is more negative than -20nm. Typically it is from -600 to -60nm. Conveniently it is from -500 to -100nm and desirably from -400 to -150nm.

[0034] Compared to WO 01/31394, the compensator layer(s) can be applied onto any and/or all of the four surfaces provided by the two liquid crystal constraints (be they glass or some other very low birefringence material). Further, the applied layers are not limited in thickness due to their coloration since they are transparent. Finally the materials suitable for these contiguous amorphous polymeric compensator layers are much broader than the aromatic polyimides of '394. A wide variety of amorphous, high glass transition temperature, polymeric materials are identified that will serve this purpose.

[0035] The invention is described in more detail by referring to the drawings as follows.

[0036] FIG. 1A shows a cross-sectional schematic of part of a liquid crystal display 5 including an amorphous polymeric compensator layer 50 and constraint 40 in accordance with the present invention. Also shown are the switchable liquid crystal 10, an alignment layer 20, a TFT (thin film transistor) layer 30, and a color filter array 35. The said amorphous polymeric compensator layer 50 having an out-of-plane birefringence more negative than -0.005, and the combined in-plane retardation (Re) of layers 20, 30, 35, 40 and 50 is from +20 to-20nm and the out-of-plane retardation (Rth) of layer 50 is more negative than-20nm.

[0037] FIG. 1B shows a cross-sectional schematic of part of another liquid crystal display 6 including an amorphous polymeric compensator layer 50 and constraint 40 in accordance with the present invention. Also shown are the switchable liquid crystal 10, an alignment layer 20, a TFT (thin film transistor) layer 30, and a color filter array 35. The said amorphous polymeric compensator layer 50 having an out-of-plane birefringence more negative than -0.005, and the combined in-plane retardation (Re) of layers 20, 30, 35, 40 and 50 is from +20 to-20nm and the out-of-plane retardation (Rth) of layer 50 is more negative than-20nm. In this embodiment layer 50 is on the other side of the constraint as compared to FIG 1A.

[0038] FIG 2A shows a liquid crystal display 700 having an amorphous polymeric compensator layer 200, a constraint 300 with alignment layer/TFT layer/color filter array on one side of the electrically switchable liquid crystal 600, a second constraint 400 with alignment layer/TFT layer which is on the other side of the electrically switchable liquid crystal 600, and polarizers 500 and 550. The transmission axes of polarizers 500 and 550 form a 90°± 10° angle relative to each other. The angles of their transmission axes are denoted as 45° and 135° relative to the liquid crystal display 700. However, other angles are possible depending on the kind of liquid crystal display 700 and this is obvious to those who skilled in the art.

[0039] FIG 2B shows another liquid crystal display 700 having two amorphous polymeric compensator layers 200, a constraint 300 with alignment layer/TFT layer/color filter array on one side of the electrically switchable liquid crystal 600, a second constraint 400 with alignment layer/TFT layer which is on the other side of the electrically switchable liquid crystal 600, and polarizers 500 and 550. The transmission axes of polarizers 500 and 550 form a 90°± 10° angle relative to each other. The angles of their transmission axes are denoted as 45° and 135° relative to the liquid crystal display 700. However, other angles are possible depending on the kind of liquid crystal display 700 and this is obvious to those who skilled in the art.

[0040] Among the liquid crystal cell arrangements useful with the invention are Vertically Aligned (VA) and In-Plane Switching (IPS) cells. In the Vertically Aligned arrangement, the liquid crystal display is one in which electric field is applied in a direction perpendicular to or vertical with respect to the cell plane and the direction of the liquid crystal optic axis is substantially perpendicular to or vertical with respect to the liquid crystal cell plane without an applied field. Therefore, the incoming light essentially does not see the birefringence through the liquid crystal cell. This state is termed the "dark state". In the In-plane switching arrangement, the electric field is applied in the direction of the liquid crystal plane and the direction of the liquid crystal optic axis is changed upon application of the electric field while remaining substantially within the plane of the cell.

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[0041] The present invention is further illustrated by the following nonlimiting examples of its practice.

Examples:

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[0042] The aromatic polyesters used herein can be prepared using any suitable or conventional procedure. The procedure used herein followed that outlined by P. W. Morgan in Condensation Polymers: By Interfacial and Solution Methods, Interscience, New York City, N.Y. (1965).

Example 1:

Polymer I (synthesis):

[0043] To a stirred mixture of 4,4"-hexafluoroisopropylidenediphenol (33.62 g, 0.1 mole) and triethylamine (22.3 g, 0.22 mole) in methylene chloride (200 mL) at 10° C. was added a solution of terephthaloyl chloride (10.15 g, 0.05 mole) and isophthaloyl chloride (10.15 g, 0.05 mole) in methylene chloride (100 mL). After the addition, the temperature was allowed to rise to room temperature and the solution was stirred under nitrogen for 4 hours, during which time triethylamine hydrochloride precipitated in a gelatinous form and the solution became viscous. The solution was then filtered and washed with dilute hydrochloric acid, (100 mL of 2% acid) followed three times by water (200 mL). The solution was then poured into methanol with vigorous stirring, and a white fibrous polymer precipitated. The glass transition temperature of this polymer was measured by differential scanning calorimetry to be 199° C.

Poly(4,4'-hexafluoroisopropylidene-bisphenol) terephthalate-co-isophthalate

Polymer I

[0044] A solution of the polyester (polymer I, 10% solids, 45% methylethylketone, 45%toluene) was coated onto a TAC web. This included the steps of unrolling the TAC web, coating the polymer solution (using a slot hopper), and applying sufficient drying (85°C) to remove the majority of the solvents. These steps occurred in a roll to roll, continuous process. Spin coating and other coating methods such as spray application could also be used. Optically clear films of the TAC/polyester structure were produced with the following optical properties. Re, Rth and the second layer thickness were measured with an ellipsometer (model M2000V, J.A. Woollam Co.) at 550nm wavelength.

TABLE I

77.022					
First Layer	SecondLayer: Polymer I Layer thickness (μm	Combined Re, In- Plane Retardation (nm)	Combined Rth, Out-of-Plane Retardation (nm)		
80μm TAC	0	3	-58		
80μm TAC	2.8	3	-84		
80μm TAC	5.6	3	-104		

Example 2:

[0045] Polymer II was similarly prepared using terephthaloyl chloride and 4,4'-(hexahydro-4,7-methanoindan-5-yli-

dene) bisphenol. The glass transition temperature of this polymer was measured by differential scanning calorimetry to be 289° C

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Poly(4,4'-hexahydro-4,7-methanoindan-5-ylidene bisphenol) terephthalate.

Polymer II

[0046] When polymer II is spun cast onto a glass substrate (10% solids in dichloroethane), it shows the following optical retardations. Re, Rth and the polymer II layer thickness are measured with an ellipsometer (model M2000V, J. A. Woollam Co.) at 550nm wavelength.

TABLE II

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Polymer II Layer thickness (μm)	Re, In-Plane Retardation (nm)	Rth, Out-of-Plane Retardation (nm)	
3.4	0.2	-74	

Example 3:

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[0047] Polymer III was similarly prepared using terephtaloyl chloride, isophthaloyl chloride and 4,4'-isopropylidene-2,2',6,6'-tetrachlorobisphenol. The glass transition temperature of this polymer was measured by differential scanning calorimetry to be 250° C.

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Poly(4,4'-isopropylidene-2,2'6,6'-tetrachlorobisphenol) terephthalate-co-isophthalate

Polymer III

[0048] When polymer III is spun cast onto glass (10% solids in dichloroethane), it shows the following optical retardations. Re, Rth and the polymer III layer thickness are measured with an ellipsometer (model M2000V, J.A. Woollam Co.) at 550nm wavelength.

TABLE III

Polymer III Layer thickness (μm)	Re, In Plane Retardation (nm)	Rth, Out of Plane Retardation (nm)
2.8	0.8	-66

Example 4:

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Polymer IV (synthesis):

[0049] To a stirred mixture of 4,4'-hexafluoroisopropylidenediphenol (23.53 g, 0.07 mole), 4,4'-(2-norbornylidene) bisphenol (8.4 g, 0.03 mole) and triethylamine (22.3 g, 0.22 mole) in methyl ethyl ketone (100 mL) at 10° C. was added a solution of terephthaloyl chloride (19.29 g, 0.095 mole) and isophthaloyl chloride (1.02 g, 0.005 mole) in methyl ethyl ketone (60 mL). After the addition, the temperature was allowed to rise to room temperature and the solution was stirred under nitrogen for 4 hours, during which time triethylamine hydrochloride precipitated in a gelatinous form and the solution became viscous. The solution was then diluted with toluene (160 mL) and washed with dilute hydrochloric acid, (200 mL of 2% acid) followed three times by water (200 mL). The solution was then poured into ethanol with vigorous stirring, and a white bead like polymer precipitated, collected and dried at 50° C. under vacuum for 24 hours. The glass transition temperature of this polymer was measured by differential scanning calorimetry to be 270° C.

Poly(4,4'-hexafluoroisopropylidene-bisphenol-co-4,4'-(2-norbornylidene) bisphenol) terephthalate-co-isophthalate.

Polymer IV

[0050] When polymer IV is spun cast onto glass (10% solids in 50% propylacetate 50% toluene), it shows the following optical retardations. Re, Rth and the polymer III layer thickness are measured with an ellipsometer (model M2000V, J.A. Woollam Co.) at 550nm wavelength.

TABLE IV

Polymer IV Layer thickness (µm)	Re, In Plane Retardation (nm)	Rth, Out of Plane Retardation (nm)	
5.9	0.2	-221	

[0051] Other specific polymers that could be used include:

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Poly(4,4'-hexafluoroisopropylidene)-bisphenol-co-(2-norbornylidene)-bisphenol terephthalate. (60/40).

[0052]

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Poly(4,4'-hexahydro-4,7-methanoindan-5-ylidene)-bisphenol-co-(4,4'-isopropylidene-2,2',6,6'-tetrabromo)-bisphenol terephthalate. (50/50)

[0053]

Poly(4,4'-isopropylidene-bisphenol-co-4,4'-(2-norbornylidene) bisphenol) terephthalate-co-isophthalate

[0054] A series of polymers were analyzed for their glass transition temperatures and out of plane birefringence values. It was found that the more desirable polymers for this invention had glass transition temperatures above 180°C. Those with lower glass transition temperatures were found to generally have birefringence values less negative than

-0.005.

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[0055] Embodiments of the invention also include those wherein:

one or more compensator layers have a combined thickness of from 0.1 to 20 micrometers, typically from 1.0 to 10 micrometers, and suitably from 2 to 8 micrometers;

the Rth of the one or more compensator layers is from -60 to -600nm, typically from -150 to -500nm;

the non-visible chromophore group includes a carbonyl, amide, imide, ester, carbonate, phenyl, naphthyl, biphenyl, bisphenol, or thiophene group;

the cell constraints are glass;

the surface is an exterior surface of the cell constraint;

a liquid crystal display comprises the liquid crystal cell of the invention; and

a process for forming a liquid crystal cell display compriesg coating the polymer of claim 1 in a solvent onto a surface of a constraint of the cell.

15 [0056] The entire contents of the patents and other publications referred to in this specification are incorporated herein by reference.

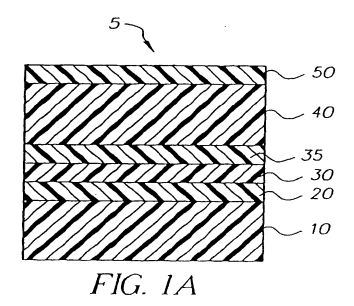
Claims

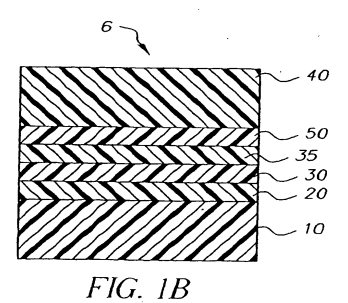
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- A liquid crystal cell having contiguous to a surface of a constraint thereof one or more compensator layers, each containing a transparent amorphous polymeric birefringent material having an out-of plane birefringence more negative than -0.005.
- 25 2. The cell of claim 1 bearing more than one of said polymeric compensator layers.
 - 3. The cell of claim 1 comprising compensator layers containing selected polymeric materials having sufficient thickness so that the overall in-plane retardation (Re) of all of the compensator layers of the liquid crystal cell is from +20 to -20nm and the out-of-plane retardation (Rth) of at least one of the compensator layers is more negative than -20nm.
 - The cell of claim 1 wherein the one or more compensator layers have a combined thickness of less than 30 micrometers.
- The cell of claim 1 wherein the out-of-plane retardation (Rth) of the one or more compensator layers is -60nm or more negative.
 - 6. The cell of claim 1 wherein the one or more compensator layers comprise a polymer containing in the backbone a non-visible chromophore group and having a Tg above 180°C.
 - The cell of claim 1 wherein the compensator layer comprises a polymer containing in the backbone a carbonyl or an aromatic group.
- 8. The cell of claim 1 wherein the compensator layer comprises 1) poly(4,4'-hexafluoroisopropylidene-bisphenol) terephthalate-co-isophthalate, 2) poly(4,4'-hexahydro-4,7-methanoindan-5-ylidene bisphenol) terephthalate, or 3) poly(4,4'-isopropylidene-2,2'6,6'-tetrachlorobisphenol) terephthalate-co-isophthalate, 4) poly(4,4'-hexafluoroisopropylidene)-bisphenol-co-(2-norbomylidene)-bisphenol terephthalate; 5) poly(4,4'-hexahydro-4,7-methanoindan-5-ylidene)-bisphenol-co-(4,4'-isopropylidene-2,2',6,6'-tetrabromo)-bisphenol terephthalate; or 6) poly(4,4'-isopropylidene-bisphenol-co-4,4'-(2-norbornylidene) bisphenol) terephthalate-co-isophthalate, or copolymers thereof.
 - 9. The compensator of claim 1 wherein the second layer comprises poly(4,4'-hexafluoroisopropylidene-bisphenol-co-4,4'-(2-norbornylidene) bisphenol) terephthalate-co-isophthalate, or copolymers thereof.
- 10. The cell of claim 1 wherein the compensator layer comprises a polymer containing in the backbone a non-visible chromophore group that does not contain a chromophore off of the backbone.
 - 11. The cell of claim 1 wherein the liquid crystal cell is a vertically aligned cell or a twisted nematic cell.

12. The cell of claim 1 wherein the liquid crystal cell is In-Plane-Switchable.





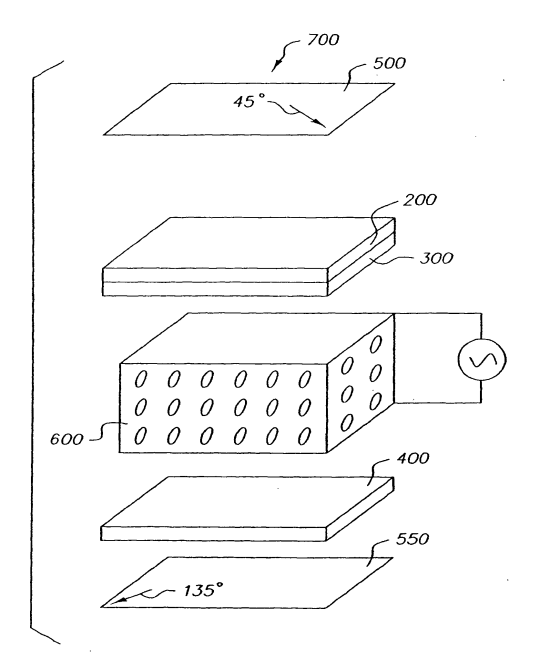


FIG. 2A

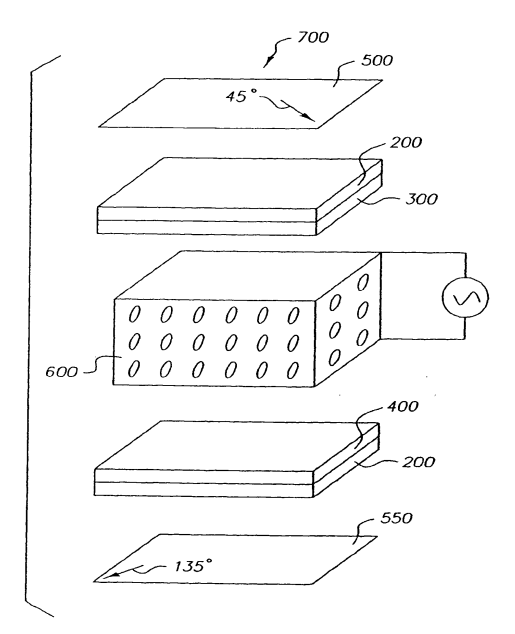


FIG. 2B



EUROPEAN SEARCH REPORT

Application Number

EP 03 07 7292

^- <u>1</u> -	Citation of document with indicati	on, where appropriate,	Relevant	CLASSIFICATION OF THE
Category	of relevant passages		to claim	APPLICATION (Int.Cl.7)
Х	US 6 380 996 B1 (ITO Y	OJI ET AL)	1-5,7,	G02F1/13363
	30 April 2002 (2002-04	-30)	10-12	C08L67/03
Y	* column 1, line 12 - * column 4, line 39 -	line 16 *	6	C08G63/193
	* column 4, line 39 -	line 46 *		
	* column 80, line 25 -	column 81, line 50		
		-		
X	US 5 699 136 A (ARAKAWA 16 December 1997 (1997)		1,11,12	
	* column 1, line 10 -	line 16 *		
	* column 4, line 13 -			
	* column 28, line 12 -	line 24 *		
	* table 3 *			
Υ	US 5 472 635 A (ITOH H	- IROYUKI ET AL)	6	
	5 December 1995 (1995-	12-05)		
	* column 2, line 10 -	line 20 *		
	* column 11, line 42 - * table 1 *	Time 52 *		
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